FIGURE 1

FIG. 1a

FIG. 1b

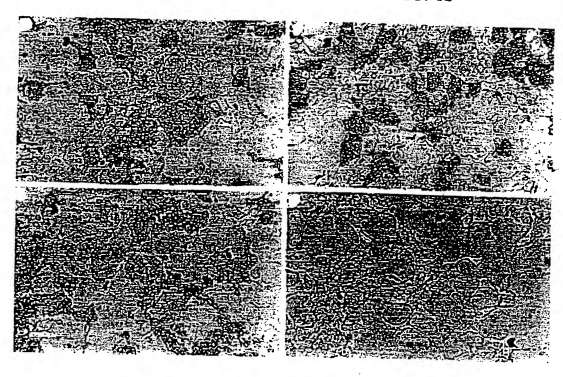
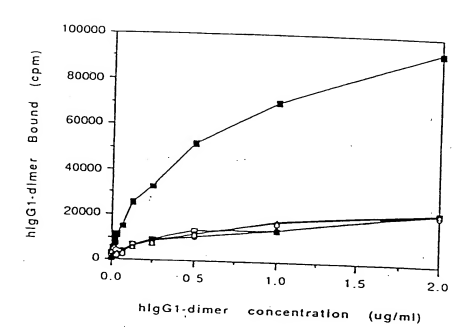
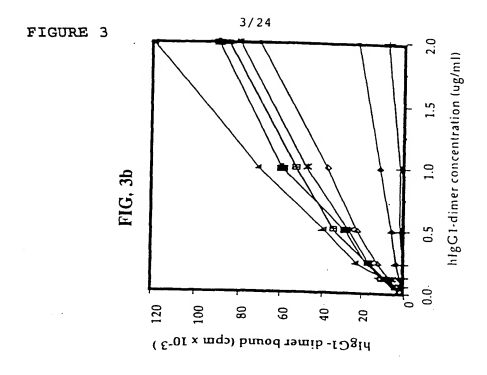
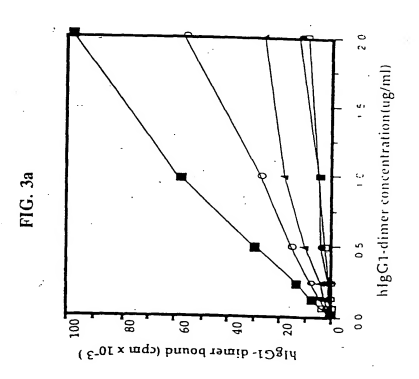


FIG. 1c

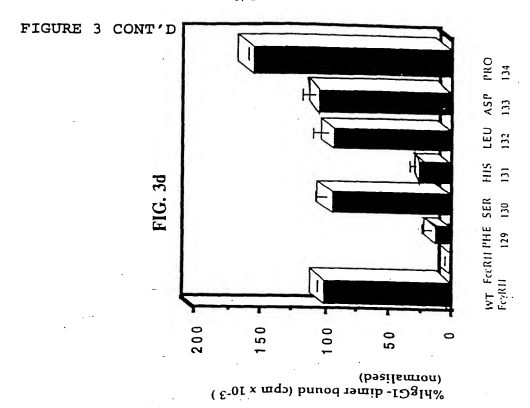
FIG. 1d

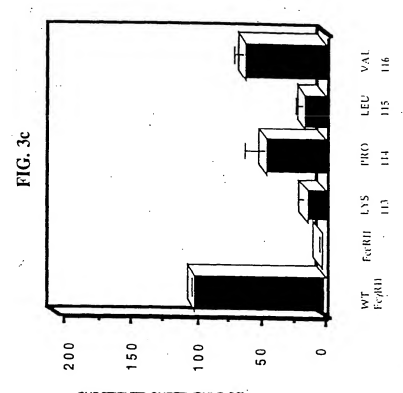






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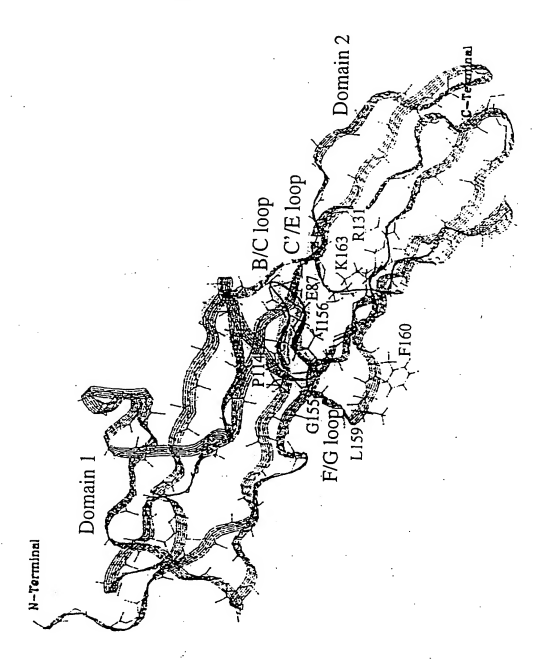


FIGURE 5

huFceRIa chain C'-E, F-G loop Ala mutant oligo nucleotides

F-G loop

5

e154 (Lys)
Ala

EFG-01 + EFG-02

EFG-01: 5' CTGTACGGGCGCAGTGTGGCAGC 3'

EFG-02:

5' GCTGCCACACTGCGCCCGTACAG 3'

10 e155 (Val) • Ala

EFG-03+EFG-04

EFG-03:

5' GTACCGGCAAAGCATGGCAGCTGG 3'

.EFG-04:

5' CCAGCTGCCATGCTTTGCCCGTAC 3'

e156 (Trp)

Ala

EFG-05+EFG-06

15 EFG-05:

5' GGGCAAAGTGGCACAGCTGGAC 3'

EFG-06:

5' GTCCAGCTGTGCCACTTTGCCC 3'

e157 (Gln)
Ala

EFG-07+EFG-08

EFG-07:

5' GCAAAGTGTGGGCACTGGACTATG 3'

20 EFG-08:

5' CATAGTCCAGTGCCCACACTTTGC 3'

e158 (Leu)
Ala

EFG-09+EFG-10

EFG-09:

5' GTGTGGCAGGCAGACTATGAGTC 3'

EGG-10:

5' GACTCATAGTCTGCCTGCCACAC 3'

25

e159 (Asp) O Ala

EFG-11+EFG-12

EFG-11:

5' GTGGCAGCTGGCATATGAGTCTG 3'

EFG-12:

5' CAGACTCATATGCCAGCTGCCAC 3'

30 e160 (Tyr)

Ala

EFG-13+EFG-14

EFG-13:

5' GCAGCTGGACGCAGAGTCTGAGC 3'

EFG-14:

5' GCTCAGACTCTGCGTCCAGCTGC 3'

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Figure 5 cont.

el61 (Glu) ® Ala

EFG-07+EFG-08

EFG-15:

5' GCTGGACTATGCATCTGAGCCCC 3'

EFG-16:

5' GGGGCTCAGATGCATAGTCCAGC 3'

5 C'-E loop

el29 (Tyr) O Ala

EBS-01 + EBS-02

EBS-01:

5' GCTCTCAAGGCATGGTATGAGAAC 3'

EBS-02:

5' GTTCTCATACCATGCCTTGAGAGC 3'

10

e130 (Trp)

Ala

EBS-03+EBS-04

EBS-03:

5' CTCAAGTACGCATATGAGAACCAC 3'

EBS-04:

5' GTGGTTCTCATATGCGTACTTGAG 3'

15 e131 (Tyτ) ⁹ Ala

EBS-01+EBS-02

EBS-05:

5' CAAGTACTGGGCAGAGAACCAC 3'

EBS-06:

5' GTGGTTCTCTGCCCAGTACTTG 3'

e132 (Glu)
Ala

EBS-07+EBS-08

20 EBS-07:

5' GTACTGGTATGCAAACCACAACATC 3'

EBS-08:

5' GATGTTGTGGTTTGCATACCAGTAC 3'

e133 (Asn)
Ala

EBS-09+EBS-10

EBS-09:

5' CTGGTATGAGGCACACACATCTCC 3'

25 EBS-10:

5' GGAGATGTTGTGTGCCTCATACCAG 3'

e134 (His) Ala

EBS-11+EBS-12

EBS-11:

5' GGTATGAGAACGCAAACATCTCCATTAC 3'

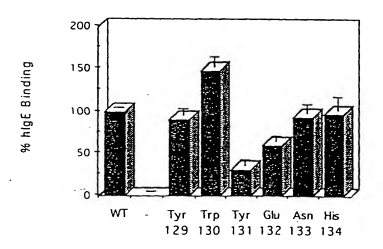
EBS-12:

5' GTAATGGAGATGTTTGCGTTCTCATACC 3'

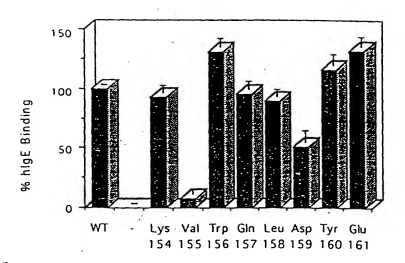
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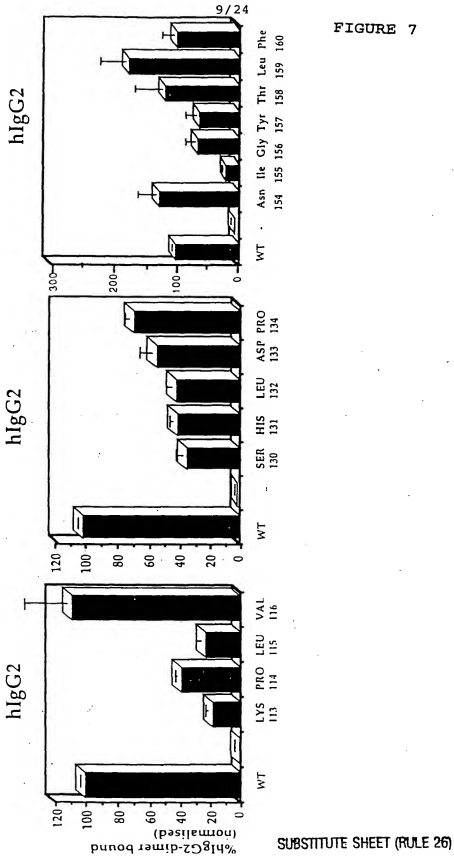
FIGURE 6

Effect of mutation of the IgE receptor (Fc ϵ RI) on the binding of IgE C'-E LOOP .









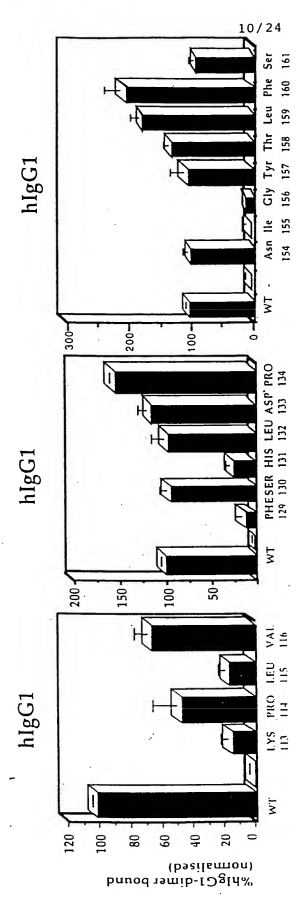


FIGURE 7 CONT'D..

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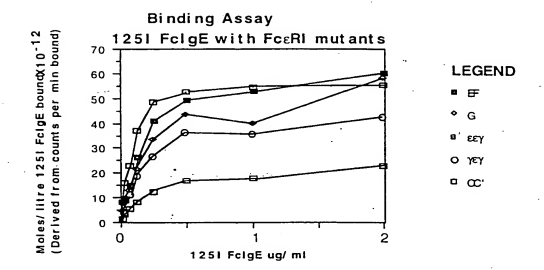
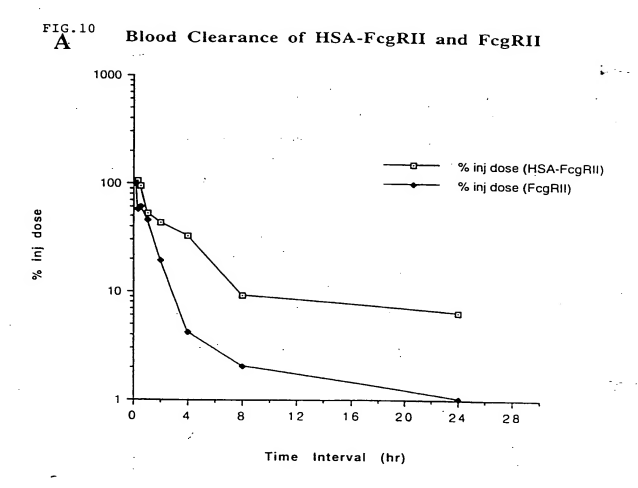


FIGURE 9

my1 y2a y2b y3 hagg 1302 8·2 1v·3 8·7

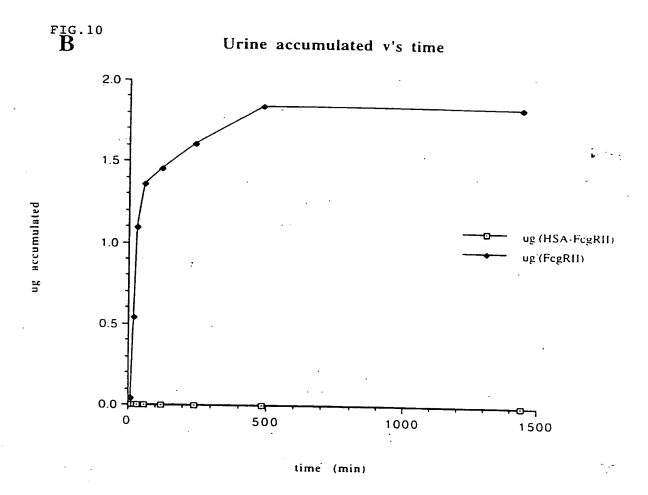
FIGURE 10



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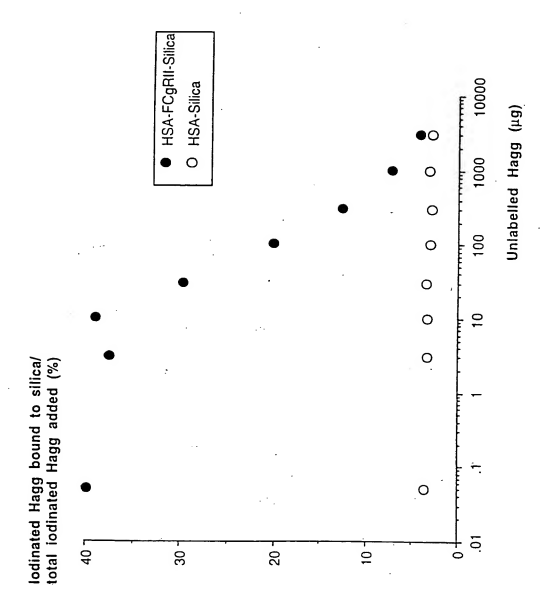
FIGURE 10 CONT'D..



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FIGURE 11

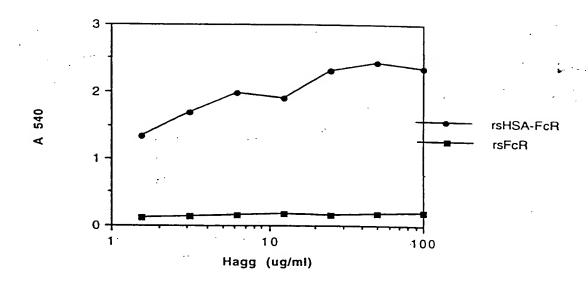


1/1									31/11									
GAT GCA	CAC	AAG	AGT	GAG	GTT	GCT	CAT	CGG			GAT	TTG	GGA	GAA	GAA	AAT	TTC	AAA
asp ala	his	lys	ser	glu	val	ala	his	arg	phe l	ys	asp	1eu	gly	glu	glu	asn	phe	lys
61/21									91/31									
GCC TTG	GTG	TTG	ATT	GCC	TTT	GCT	CAG	TAT	CTT C	AG	CAG	TGT	CCA	TTT	GAA	GAT	CAT	GTA
ala leu 121/41	val	Ieu	ıle	ala	pne	aıa	gın	суг	151/5		gin	cys	pro	pne	giu	asp	nıs	vai
AAA TTA	GTG	TAA	GAA	GTA	ACT	GAA	TTT	GCA		_	TGT	GTT	GCT	GAT	GAG	TCA	GCT	GAA
lys leu																		
181/61									211/7									40.
AAT TGT asn cys	GAC	AAA	TCA	CTT	CAT	ACC	CTT	TTT	GGA G	iAC	AAA	TTA	TGC	ACA	GTT	GCA	ACT	CTT
241/81	asp	ıys	ser	Ieu	nis	cnr	Ten	pne	271/9		TAR	reu	cys	CIII	Val	ala	CHI	Ten
CGT GAA	ACC	TAT	GGT	GAA	ATG	GCT	GAC	TGC	TGT G	CA	AAA	CAA	GAA	CCT	GAG	AGA	AAT	GAA
arg glu	thr	tyr	gly	glu	met	ala	asp	cys	cys a	la	lys	gln	glu	pro	glu	arg	asn	glu
301/101					~ · m	a.a		CC3	331/1		ccc	CCX	mm-	CTC	202	CCA	C. C.	C C C C C C C C C C C C C C C C C C C
TGC TTC	TTG	CAA	CAC	AAA	GAT	GAC	AAC	DTO	asn 1	eu	pro	ara	leu	val	ara	DYO	GAG	GTT val
361/121		gin	1112	Lys	αэр	asp	u	P	391/1		F	5				220	914	•41
GAT GTG	ATG	TGC	ACT	GCT	TTT	CAT	GAC	AAT	GAA C	GAG	ACA	TTT	TTG	AAA	AAA	TAC	TTA	TAT
asp val		cys	thr	ala	phe	his	asp	asn			thr	phe	leu	lys	lys	tyr	leu	tyr
421/141 GAA ATT	CCC		202	C.M	CCT	TO C	THE	ጥልጥ	451/1 GCC C		GAA	CTC	CONT	שייר	ערעיער	CCT	מממ	ACC.
glu ile	ala	AGA	ara	his	nra	tvr	phe	tvr	ala r	oro	alu	leu	leu	phe	phe	ala	lvs	arg
481/161									511/1	171								
TAT AAA	GCT	GCT	TTT	ACA	GAA	TGT	TGC	CAA	GCT C	CT	GAT	AAA	GCT	GCC	TGC	CTG	TTG	CCA
tyr lys		ala	phe	thr	glu	cys	cys	gln			asp	lys	ala	ala	cys	leu	leu	pro
541/181 AAG CTC	CAM	C 3 3	Cutu	ccc	СМТ	CAA	ccc	AAG	571/1 GCT 1		ጥርጥ	CCC	222	CAG	AGA	CTC	DAA	ጥርጥ
lys leu	asp	alu	leu	arg	asp.	glu	alv	lys	ala s	ser	ser	ala	lys	gln	arg	leu	lys	cys
601/201									631/2	211								
GCC AGT	CTC	CAA	AAA	TTT	GGA	GAA	AGA	GCT	TTC A	AAA	GCA	TGG	GCA	GTA	GCT	CGC	CTG	AGC
ala ser		gln	lys	phe	gly	glu	arg	ala	phe 1		ala	trp	ala	val	ala	arg	leu	ser
661/221 CAG AGA	ጥጥጥ	CCC	222	CCT	GAG	ጥጥጥ	GCA	GAA			AAG	TTA	GTG	ACA	GAT	СТТ	ACC	AAA
gln arg	phe	pro	lvs	ala	glu	phe	ala	glu	val s	șer	lys	leu	val	thr	asp	leu	thr	lys
721/241									751/3	251								
GTC CAC	ACG	GAA	TGC	TGC	CAT	GGA	GAT	CTG	CTT	GAA	TGT	GCT	GAT	GAC	AGG	GCG	GAC	CTT
val his 781/261		glu	cys	cys	his	gry	asp	reu	811/2	91U 271	cys	ala	asp	asp	arg	ala	asp	reu
GCC AAC	TAT	ATC	TGT	GAA	AAT	CAA	GAT	TCG			AGT	AAA	CTG	AAG	GAA	TGC	TGT	GAA
ala lys	tyr	ile	cys	glu	asn	gln	asp	ser	ile	ser	ser	lys	leu	lys	glu	cys	cys	glu
841/281									871/	291								
AAA CCT	CTC	TTG	GAA	AAA	TCC	CAC	TGC	ATT	GCC	GAA	GTG	GAA	TAA	GAT	GAG	ATG	CCL	GC".
1ys pro		ı ıeu	gin	ıys	ser	nis	cys	116	931/			gra	a 511	asp	910	mec	pro	ara
GAC TTO	CCI	TCA	TTA	GCT	GCT	GAT	TTT	GTT	GAA .	AGT	AAG	GAT	GTT	TGC	AAA	AAC	TAT	GCT
asp lev	pro	ser	leu	ala	ala	asp	phe	val	glu	ser	lys	asp	val	суs	lys	asn	tyr	ala
961/321	l								991/								- C-OM	Chmir
GAG GC	A AAC	GAI	GTC	TTC	CTG	GGC	ATG	TTT	THE	TAT	GAA	TAT	GCA	AGA arn	AGG	his	nro	GAI;
1021/3		asp) vai	. pne	. reu	gry	mer	. pne	1051	/35	1		414	9	4.9		pro	005
TAC TC	r GTC	GTO	CTC	CTC	CTG	AGA	CTI	GCC	AAG	ACA	TAT	' GAA	ACC	ACT	CTA	GAG	AAG	TGC
tyr se	r val	l val	leu	lev	ı leu	arç	leu	ala	lys	thr	tyr	glu	thr	thr	leu	glu	lys	cys
1081/3	51								1111			mmc	C 2 /		- Chestra		CC	י כייי
TGT GCC	GC'	r GCA	GAT	CCI	CAT	GA/	TGC	. IAI	ala	lve	val	phe	. GAT	. GAA	. il.	lve	pro	leu
1141/3	81						_		1171	./39	1							
GTG GA	A GA	G CCI	CAC	S AAT	ATT 7	ATO	XAA	CAA	TAA	TGT	GAC	CTI	TTT	GAC	CAC	CTI	GGA	GAG
val gl	u gli	u pro	glr	n asr	ı lev	ile	a lys	gln	asn	cys	glu	lev	phe	glı	glr	ı lev	gly	glu
1201/4	01								1231			2 332	Cura			CTC	ב קרר א	АСТ
TAC AA	A TT	CAC	AA?	r GCC	CTA	177	4 GT"	L GGT	TAC	thr	· lve	· AAA	val	י בכנ מיים	. CAA	i val	ser	thr
1261/4		e gri	ı ası	1 415	ı ret	. 161	. va.		1291	/43	1	1 -						-
, 1	~ _																	

FIGURE 12 CONT'D...

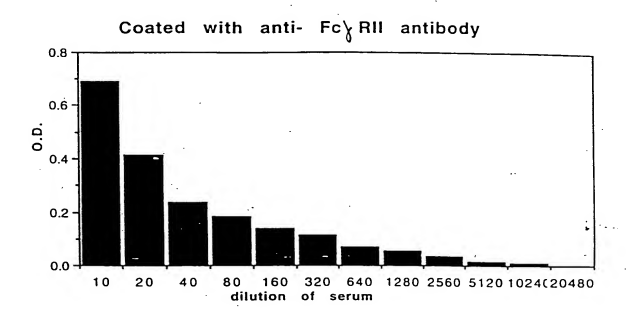
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pro thr leu val glu val ser arg asn leu gly lys val gly ser lys cys cys lys his
1321/441
                                        1351/451
CCT GAA GCA AAA AGA ATG CCC TGT GCA GAA GAC TAT CTA TCC GTG GTC CTG AAC CAG TTA
pro glu ala lys arg met pro cys ala glu asp tyr leu ser val val leu asn gln leu
1381/461
                                        1411/471
TGT GTG TTG CAT GAG AAA ACG CCA GTA AGT GAC AGA GTC ACA AAA TGC TGC ACA GAA TCC
cys val leu his glu lys thr pro val ser asp arg val thr lys cys cys thr qlu ser
                                        1471/491
1441/481
TTG GTG AAC AGG CGA CCA TGC TTT TCA GCT CTG GAA GTC GAT GAA ACA TAC GTT CCC AAA
leu val asn arg arg pro cys phe ser ala leu glu val asp glu thr tyr val pro lys
1501/501
                                        1531/511
GAG TTT AAT GCT GAA ACA TTC ACC TTC CAT GCA GAT ATA TGC ACA CTT TCT GAG AAG GAG
glu phe asn ala glu thr phe thr phe his ala asp ile cys thr leu ser glu lys qlu
                                        1591/531
1561/521
AGA CAA ATC AAG AAA CAA ACT GCA CTT GTT GAG CTC GTG AAA CAC AAG CCC AAG GCA ACA
arg gln ile lys lys gln thr ala leu val glu leu val lys his lys pro lys ala thr
                                        1651/551
1621/541
AAA GAG CAA CTG AAA GCT GTT ATG GAT GAT TTC GCA GCT TTT GTA GAG AAG TGC TGC AAG
lys glu gln leu lys ala val met asp asp phe ala ala phe val glu lys cys cys lys
1681/561
                                        1711/571
GCT GAC GAT AAG AAG ACC TGC TTT GCC GAG GAG GGT AAA AAA CTT GTT GCT GCA AGT CAA
ala asp asp lys lys thr cys phe ala glu glu gly lys lys leu val ala ala ser gln
                                         1771/591
1741/581
GCT GCC TTA GGC GCT CCC CCA AAG GCT GTG CTG AAA CTT GAG CCC CCG TGG ATC AAC GTG
ala ala leu gly ala pro pro lys ala val leu lys leu glu pro pro trp ile asn val
                                        1831/611
1801/601
CTC CAG GAG GAC TCT GTG ACT CTG ACA TGC CAG GGG GCT CGC AGC CCT GAG AGC GAC TCC
leu gln glu asp ser val thr leu thr cys gln gly ala arg ser pro glu ser asp ser
                                         1891/631
1861/621
ATT CAG TGG TTC CAC AAT GGG AAT CTC ATT CCC ACC CAC ACG CAG CCC AGC TAC AGG TTC
ile gln trp phe his asn gly asn leu ile pro thr his thr gln pro ser tyr arg phe
1921/641
                                         1951/651
AAG GCC AAC AAC AAT GAC AGC GGG GAG TAC ACG TGC CAG ACT GGC CAG ACC AGC CTC AGC
lys ala asn asn asp ser gly glu tyr thr cys gln thr gly gln thr ser leu ser
                                         2011/671
1981/661
GAC CCT GTG CAT CTG ACT GTG CTT TCC GAA TGG CTG GTG CTC CAG ACC CCT CAC CTG GAG
asp pro val his leu thr val leu ser glu trp leu val leu gln thr pro his leu glu
2041/681
                                         2071/691
TTC CAG GAG GGA GAA ACC ATC ATG CTG AGG TGC CAC AGC TGG AAG GAC AAG CCT CTG GTC
phe gln glu gly glu thr ile met leu arg cys his ser trp lys asp lys pro leu val
2101/701
                                         2131/711
AAG GTC ACA TTC TTC CAG AAT GGA AAA TCC CAG AAA TTC TCC CAT TTG GAT CCC ACC TTC
lys val thr phe phe gln asn gly lys ser gln lys phe ser his leu asp pro thr phe
                                         2191/731
2161/721
TCC ATC CCA CAA GCA AAC CAC AGT CAC AGT GGT GAT TAC CAC TGC ACA GGA AAC ATA GGC
ser ile pro gln ala asn his ser his ser gly asp tyr his cys thr gly asn ile gly
2221/741
                                         2251/751
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tyr thr leu phe ser ser lys pro val thr ile thr val gln AMB
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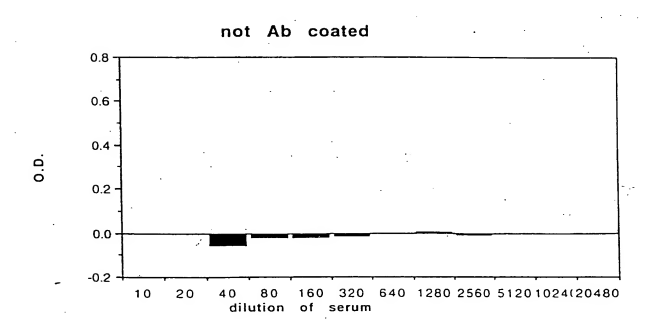
FIGURE 13



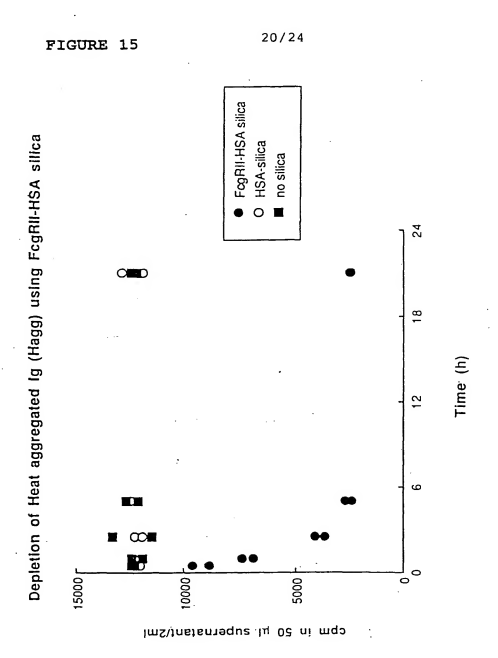
19/24

FIGURE 14

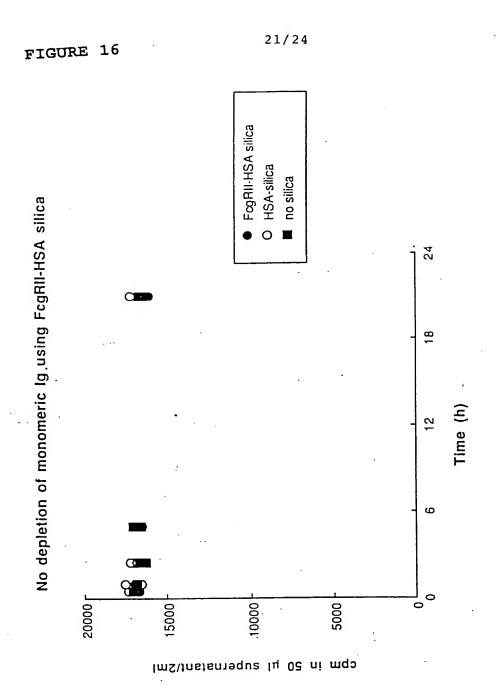




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SUBSTITUTE SHEET (Rule 26)



SUBSTITUTE SHEET (Rule 26)

Functional soluble FcyRII detected binding to hagg using the MoAb 8.2

Titration of rsFcgRII from various sources

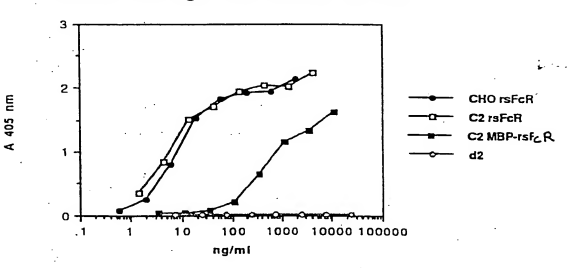
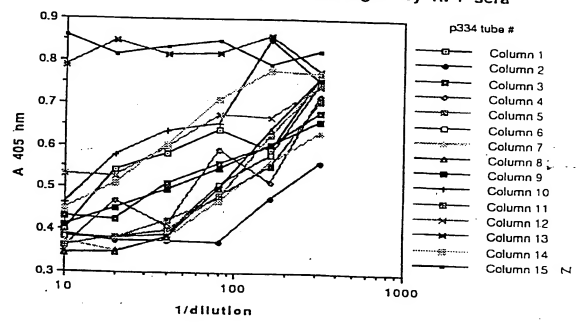


FIGURE 18

Human serum containing rhuematoid factors, but not normal human sera, inhibits the binding of HRP conjugated to rsHSA-FcyRII to hagg.

Inhibition of HRP labelled rsHSA-FcgRII by RF+ sera



Peptoid	A 405 nm	Peptoid	A 405 nm
name	(X 1000)	name	(X 1000)
TC,1 AD,1 DC,1 SC,1. AC,1 RC,1 RC,1 UC,1 LC,1 FC,1 JC,1 HC,1 QC,1 GC,1 CU,1 MC,1 EC,1 DG,1 IC,1 CF,1 CV,1 VC,1	308.5 317 319 319 320 323.5 329.5 329.5 330.5 332.5 335 343 345.5 346.5 356 357 358 359.5 363 363 364 364 364	NB,1 BD,1 OA,1 DJ,1 LG,1 VF,1 OG,1 DS,1 IF,1 RA,1 DL,1 QB,1 QF,1 EU,1 JE,1 DQ,1 AF,1 SB,1 HA,1 IG,1 HB,1 DO,1 RB 1 Background Maximum	578.5 580 582.5 586 586.5 586.5 587 588 588 589.5 591 592 592 592 592 592 592 594 594 594 594,5 596 596.5 596.5 597 304 619